



Myalgic Encephalomyelitis/Chronic Fatigue Syndrome



MAY 10, 2024

CDC's website is being modified to comply with President Trump's Executive Orders.

ME/CFS Basics

KEY POINTS

- Myalgic encephalomyelitis/chronic fatigue syndrome (ME/CFS) is a serious and often long-lasting illness that keeps people from doing their usual activities.
- It makes physical and mental exertion difficult.
- Symptoms include trouble thinking, severe tiredness and other symptoms.
- There is no known cause or cure. Care usually means treating the symptoms that most affect a person's life.



MORE INFORMATION

- For Everyone
- Health Care Providers

Overview

ME/CFS is a biological illness that affects many body parts. It causes severe fatigue not improved by rest, problems thinking and sleeping, dizziness, pain, and many other symptoms.

People with ME/CFS may not look sick but can't do their normal activities. ME/CFS may get worse after they do any activity -- physical or mental. This symptom is called post-exertional malaise (PEM). After they exert themselves, they may need to stay in bed for an extended time. About 1 in 4 people with ME/CFS are confined to bed at some point in their illness.

Did you know?



It's estimated that up to 3.3 million people in the United States suffer from ME/CFS. More than 9 in 10 people with ME/CFS have not been diagnosed by a doctor. ME/CFS costs the U.S. economy about \$18 to \$51 billion annually in medical costs and lost income.

It is difficult to diagnose because there is no specific test for ME/CFS. One reason that people with ME/CFS have not been diagnosed may be limited access to healthcare. Another reason may be a lack of healthcare providers who know about ME/CFS and how to recognize it.

Anyone can get ME/CFS. It doesn't matter what age, gender or ethnicity they are. However, some groups are more affected than others.

Signs and symptoms

People with ME/CFS are not able to function the same way they did before they became ill. They may not always be able to do daily tasks like showering or cooking a meal.

ME/CFS often makes it hard to keep a job, go to school, and take part in family and social life. ME/CFS can last for years and sometimes leads to serious disability.

ME/CFS is most common in people in middle age but also affects kids, teens and elderly people. Among adults, women are affected more often than men. Among children, teens are more affected than younger kids.

People who are white are diagnosed more often than other races and ethnicities. But many people with ME/CFS have not been diagnosed, especially among people from racial and ethnic minority groups.

Diagnosis

Symptoms of ME/CFS may appear similar to many other illnesses. There is no test to confirm ME/CFS. This makes it difficult to diagnose. The illness can also be unpredictable. Symptoms may come and go, or there may be changes in how severe symptoms are over time.

A doctor should be able to distinguish ME/CFS from other illnesses by doing a thorough medical exam. This includes asking many questions about the patient's health history and current illness. They should also ask about how often symptoms occur, how bad they are, and how long they have lasted. It is also important for doctors to talk with patients about how the symptoms affect their lives.

Treatment and recovery

There is no cure or approved treatment for ME/CFS. But some symptoms can be treated or managed. Treating these symptoms might provide relief for some people with ME/CFS but not others. Other strategies, like learning new ways to manage activity, can also help.

People with ME/CFS, their families, and healthcare providers need to work together to decide which symptom causes the most problems. This should be treated first after discussing possible benefits and harms of any treatment plans, including medicines and other therapies.

Healthcare providers need to support their patients' families as they come to understand how to live with this illness. Providers and families should remember that this process is likely to be hard on people with ME/CFS.

Resources

Could You Have ME/CFS? (Myalgic Encephalomyelitis/Chronic Fatigue Syndrome)

ME/CFS is a complex illness and symptoms of ME/CFS may seem similar to many other illnesses. ME/CFS requires **three** symptoms:

1 Not being able to participate in routine activities that were possible before becoming ill, such as work, school, social life, and/or personal life, that:

- Lasts for more than **6 months**
- Is accompanied by **fatigue** that is:
 - Often serious
 - Just started (not lifelong)
 - Not the result of ongoing activities
 - Not from more than usual effort
 - Not made better by rest

2 Post-exertional malaise (PEM). Worsening of symptoms after physical, mental, or emotional effort that would not have caused a problem before the illness. This is sometimes referred to as "crashing" by people with ME/CFS.

3 Unrefreshing sleep. People with ME/CFS may not feel better even after a full night of sleep (e.g., feeling just as tired upon waking up as before going to bed).


In addition, **at least one** of the following symptoms is also required:

Impaired memory or ability to concentrate. People with ME/CFS may have trouble remembering, learning new things, concentrating, or making decisions.

Orthostatic intolerance (symptoms that occur when standing upright). People with ME/CFS may feel lightheaded or dizzy when standing upright and may even faint.

The list of key symptoms is drawn from an Institute of Medicine (IOM) report by an expert committee of the National Academies of Sciences, Engineering, and Medicine and published in 2015: *Beyond Myalgic Encephalomyelitis/Chronic Fatigue Syndrome: Redefining an Illness*. You may experience some additional symptoms.

Only a healthcare provider can diagnose ME/CFS. A healthcare provider will ask about how often your symptoms occur and how much they affect you. Sometimes you may need to make more than one visit to a healthcare provider before being diagnosed. While not all healthcare providers are familiar with diagnosing ME/CFS, resources are available to help them make a diagnosis.



Centers for Disease Control and Prevention
National Center for Emerging and Zoonotic Infectious Diseases

For more information on ME/CFS, please visit www.cdc.gov/me-cfs.

Patient Toolkit

ME/CFS educational tools to help patients and their families manage their healthcare visits.

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Living with ME/CFS

Patients and their families need support as they come to understand how to live with ME/CFS.

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SOURCES

CONTENT SOURCE:

National Center for Emerging and Zoonotic Infectious Diseases (NCEZID)

SOURCES

- [Institute of Medicine \(IOM\) report](#) 
- [NCHS Data Brief, Number 488, December 2023 \(cdc.gov\)](#) 
- <https://www.frontiersin.org/articles/10.3389/fped.2019.00185/full> 